

(43) International Publication Date
14 August 2014 (14.08.2014)

(51) International Patent Classification:

C22C 32/00 (2006.01) C22C 29/12 (2006.01)
C22C 21/00 (2006.01)

(21) International Application Number:

PCT/CA2014/000102

(22) International Filing Date:

11 February 2014 (11.02.2014)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

61/763,186 11 February 2013 (11.02.2013) US

(71) Applicant: NATIONAL RESEARCH COUNCIL OF CANADA [CA/CA]; 1200 Montreal Road, Ottawa, Ontario K1A 0R6 (CA).

(72) Inventor: BOUCHARD, Dominique; 805 Saguenéens, unité 12, Chicoutimi, Quebec G7H 0A3 (CA).

(74) Agents: DAVIS, Jason, E., J. et al.; IP Portfolio Management, National Research Council of Canada, 1200 Montreal Road, Bldg. M55-Room 29, Ottawa, Ontario K1A 0R6 (CA).

(81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM,

AO, AT, AU, AZ, BA, BB, BG, BH, BN, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IR, IS, JP, KE, KG, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PA, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SA, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

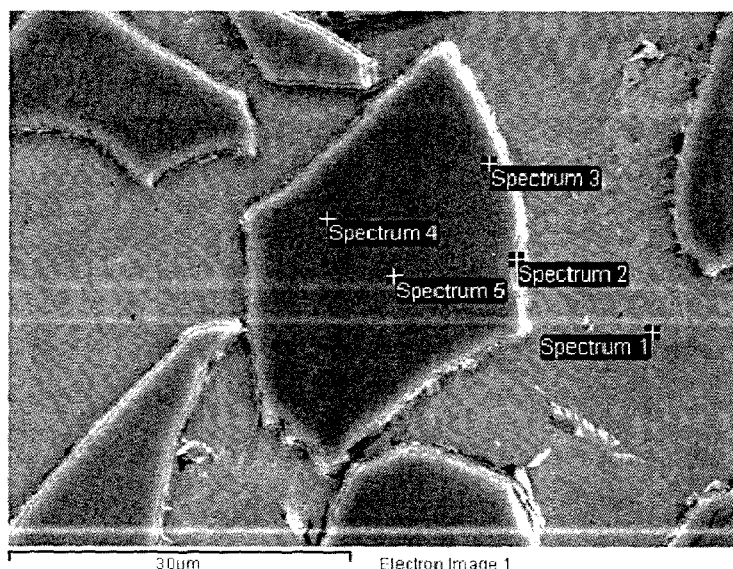
(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, RU, TJ, TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, KM, ML, MR, NE, SN, TD, TG).

Declarations under Rule 4.17:

- as to applicant's entitlement to apply for and be granted a patent (Rule 4.17(ii))
- of inventorship (Rule 4.17(iv))

[Continued on next page]

(54) Title: METAL MATRIX COMPOSITE AND METHOD OF FORMING

**FIG. 2c**

(57) Abstract: Use of Ca in metal matrix composites (MMC) allows for incorporation of small and large amounts of ceramic (e.g. rutile TiO₂) into the metal (Al, or its alloys). Calcium remains principally out of the matrix and is part of a boundary layer system that has advantages for integrity of the MMC. Between 0.005 and 10 wt.% calcium (Ca) may be included, and more than 50wt.% of rutile has been shown to be integrated. Rutile may therefore be used to reduce melt loss due to calcium from an aluminum or aluminum alloy melt.

WO 2014/121384 A1



Published:

— *with international search report (Art. 21(3))*

METAL MATRIX COMPOSITE AND METHOD OF FORMING

Field of the Invention

[0001] The present invention relates in general to metal matrix composites (MMCs) and methods of forming MMCs, and in particular to the use of calcium to improve integration of ceramics in aluminum containing metal matrices.

Background of the Invention

[0002] MMCs are a class of materials having many applications where mechanical properties such as strength, abrasion resistance, thermal resistance, or lightness are sought. MMCs are composed of a metal matrix and reinforcement. Herein the reinforcements include, and are preferably composed principally of, ceramics or cermets. There are many fabrication routes for generating MMCs, but typically a lowest cost route involves melting the metal, adding powdered ceramics or cermets, stirring, and then cooling the mixture to solidify. This production route is often called 'stir casting'. The cooling may be performed by casting the mixture, by injection molding or by extrusion using a variety of techniques known in the art.

[0003] There are problems in the art with choosing reinforcement and metal materials. Some candidates react with each other. For example, it was natural to try carbon fibers in aluminum, as both are used in the aerospace industry for their lightness and strength. However, aluminum reacts with carbon to form Al_4C_3 , which is brittle, moisture sensitive, and therefore problematic. Therefore carbon fibers are typically coated to prevent this reaction. Such coatings add cost and difficulties to the production of MMCs, and introduce other problems. The coating has to reliably passivate the carbon, on one side and present a non-reactive surface to the metal on the other.

[0004] If the reinforcement is selected (or coated) so that it does not react with the molten metal, there is still an important hurdle to producing useful MMCs: integration. The interfaces between the reinforcement and the liquid metal, when there is low affinity between the metal and reinforcement, are crucial to the strength of the material. Liquid metals and particularly aluminum typically exhibit poor wetting with reinforcement particles. In many cases this is attributable to the formation of a matrix oxide layer at the interface with the particles that hinders intimate contact. If the interfaces are not wetted, even with good mixing, and equal net forces on the reinforcements and metal, separation of the reinforcements and metal are likely, leading to a generally unwanted bulk mixture that is

heterogeneous. This heterogeneity may be exacerbated by thermal contraction during solidification, which typically affects the metal much more than the reinforcements.

[0005] The more ceramic in the mixture, the more wetting is required to produce a MMC solid that is free of voids to form monolithic, integrated materials. Generally, the smaller the sizes of the surfaces of the reinforcement, the more wetting is required for integration. This is unfortunate because it is desired to retain small reinforcement particle sizes for some applications, and a range of reinforcement to matrix ratios are frequently desired.

[0006] Thus it is known in the art to use wetting agents in liquid metal and ceramic mixtures to promote intimate contact between the powders and metal. Magnesium seems to be the preferred wetting agent. For example, [1] Chaudhury teaches a stir casting method of producing a MMC with Al as the metal, and rutile TiO_2 powders as the reinforcement. It is noted that using finer rutile particles led to a high rejection rate, and limited amounts of the powder could be retained in the melt. About 2 wt.% of magnesium was plunged into the melt to increase wettability. Even with the Mg, only 11 wt.% of TiO_2 was successfully incorporated into the melt, and a greater degree of segregation of the TiO_2 from the Al was observed at the top in comparison with the bottom of the castings, which indicates a lack of uniformity. Furthermore microvoids were observed in the particle rich zones.

[0007] According to [2] Hashim et al., addition of alloying elements can help. Excellent bonding between ceramic and molten matrix can be achieved when reactive elements are added to induce wettability. For example, addition of magnesium, calcium, titanium, or zirconium to the melt may promote wetting by reducing the surface tension of the melt, decreasing the solid-liquid interfacial energy of the melt, or inducing wettability by chemical reaction. According to [2], it has been found that magnesium has a greater effect in incorporating reinforcement particles into aluminum based melts than others that were tried, including cerium, lanthanum, zirconium, titanium, bismuth, lead, zinc, and copper. Mg successfully promotes wetting of alumina, and is thought to be suitable in aluminum with most reinforcements.

[0008] [3] Rohatgi reviews cast Al MMCs for automotive applications. It mentions that stir casting and pressure infiltration are two solidification techniques that both require mixing and wetting between the molten alloys and reinforcements. According to [3]: "High-strength, high-stiffness polycrystalline α -alumina (Al_2O_3)/Al composites have been prepared by a pressure-infiltration process. For nonwetting metals, the α - Al_2O_3 is coated with a metal by vapor deposition or by electroless plating before infiltration. Titanium-boron coatings have also been used for graphite (Gr)/Al and Al_2O_3 /Al composites. However, in terms fabricability

and cost, modification of the matrix by adding small amounts of reactive elements (e.g., Mg, Ca, Li or Na) is preferred. Alumina-reinforced aluminum composites, as well as several particle-filled MMCs, have been synthesized by adding reactive agents to the melts."

[0009] Typically MMCs produced by stir casting (as opposed to the infiltration techniques that can incorporate very large amounts of reinforcements but require a costly and time-consuming ceramic pre-form to be fabricated beforehand) are substantially limited in the amount of reinforcement they can include. So the table III of Al MMCs in [3] shows that all of the MMCs have 5-20 wt.% of reinforcements, except Lanxide, which used the pressure infiltration process, which is more expensive than the preferred stir casting technique (as expressly noted therein). It should also be noted that the very high concentrations of reinforcements in these applications are associated with significantly greater strength and modulus than the 5-20 wt.% MMCs. All of the reinforcements used were ceramic powders (except for short fibres used by Honda).

[0010] Some information can be gleaned about the effect of calcium on surface tension from work on metal foams, and the distribution of calcium oxide within foamed metal, for example from [4] Hui, and [5] Banhart. While it is not exactly clear in these two references what the effect is, it does appear to have a notable effect on the viscosity and surface tension of a foaming metal. Per [4], the surface tension of commercially pure Al, drops rapidly with the addition of 2 wt.% of Ca.

[0011] While calcium may be included in foamed metal compositions in order to control frothing, calcium is not a particularly inviting element to include in Al melts. According to [6] calcium, lithium, and sodium are elements that are regarded as impurities in many aluminum alloys. The impurities contribute to the rejection rate of aluminum sheet and bar products. Rejected products must be remelted and recast. During this process, a portion of the aluminum is lost to oxidation (melt loss). Removal of calcium, lithium, and sodium increase overall melt loss of aluminum alloys. These impurities increase the hydrogen solubility in the melt and promote the formation of porosity in aluminum castings. According to Aluminum Alloys Castings Properties, Processes and Applications Chapter 2 / 15, Section 2.5.6: Calcium is a weak aluminum-silicon eutectic modifier. It increases hydrogen solubility and is often responsible for casting porosity at trace concentration levels. Calcium greater than approximately 0.005% also adversely affects ductility in aluminium-magnesium alloys.

[0012] Accordingly there is a need for a technique for improving integration of ceramic powders into molten metal to produce MMCs that can be stir cast, for example, especially techniques that allow for the integration of a greater amount of the ceramic powders.

Summary of the Invention

[0013] While Ca may offer an essential control for the foaming of metal, and while Ca is included in several lists of possible, untried, wetting agents possibly suitable for Al for melt casting, and even though Ca is known to decrease surface tension of Al, it had not been tried, it was not obvious to work as a wetting agent, it was not obvious that working as a wetting agent, or other agent for improving integration, that it wouldn't also lead to high rejection rates of MMCs.

[0014] Applicant has unexpectedly discovered that calcium is a far better additive to promote integration of ceramics in aluminum than magnesium is, at least when the ceramic is rutile TiO_2 , or the like. In fact, the use of Ca, in small amounts, has a remarkable ability to allow for more than 50 wt. % of rutile TiO_2 into an aluminum melt with a stir casting technique. No high concentration stir-cast MMCs were previously known in the art. Anatase TiO_2 (a polymorph of TiO_2 different only from rutile in a crystal structure) was tried and it did not integrate well with the melt with equal amounts of Ca, which shows that the knowledge that Ca reduces surface tension of Al does not ensure that it would improve the integration of powders of reinforcing ceramics. The rutile polymorph is inherently more stable than the anatase, so if free energy were a guide, it would be expected that anatase would be the more likely polymorph to form a stable metal-ceramic interface. Apparently kinetic barriers are still present for the incorporation of particles even when a reduction of surface tension conducive to improved particle wetting has been achieved. Therefore, the effect of calcium additions to improve the integration of rutile in liquid aluminum cannot be explained only in terms of its role as a wetting agent. Furthermore, while Ca is a stronger oxygen scavenger than Ti or Al, it was by no means certain that Ca would be substantially confined to the oxide-containing ceramic regions of the MMC, as was found. Finally, a calcium-containing boundary system appears to form around rutile that is associated with improved integration with the Al-containing matrix.

[0015] Accordingly, a method for producing a metal matrix composite is provided, the method comprising mixing a reinforcement with an aluminum-containing molten or semisolid metal or alloy and between 0.005 and 10 wt.% calcium (Ca), wherein the reinforcement is composed of particles each having a surface bearing at least 20% of titanium oxide (TiO_2), and the TiO_2 is predominantly of crystal form other than anatase; and cooling the mixture to produce a solid metal matrix composite.

[0016] The reinforcement may be a cermet or ceramic powder including the TiO_2 , or a compound coated with the TiO_2 . The TiO_2 may be in a rutile or brookite crystal form. Rutile

TiO₂ has been proven. The mixture may consist of at least 60 wt.%, more preferably 80 wt.%, more preferably 90 wt.%, more preferably 95 wt.%, more preferably 97 wt.% of the reinforcement and molten metal. The molten or semisolid metal may be liquid aluminum of a predetermined purity.

[0017] The molten metal may include aluminum, and at least one alloying metal in liquid or semisolid form with the aluminum, other than magnesium. The molten metal may be composed of more Al than any other element by weight.

[0018] The particles may be spherical, cubic, prismatic, polyhedral, angular, amorphous, elongated, rod-like, tubular, conic, fibrous, filamentary, platelet-like, disc-like, irregular, or any combination of the above. The surfaces of the particles may be flat, or curved, smooth or rough, randomly textured or patterned, concave or convex, or any combination of the above. The particles may have a predefined distribution of dimensions, with less than 10% of the reinforcements having dimensions greater than a maximum dimension, which is less than 1 cm, and with less than 10% of the reinforcements having dimensions smaller than a minimum dimension, which is greater than 10 nm. Each surface of the typical particle may bears at least 20%, or more preferably at least 60% of TiO₂.

[0019] Cooling the mixture to produce a solid metal matrix composite may comprise: sandcasting, die casting, centrifugal casting, compocasting, thixocasting, rheocasting, thixomolding or other semisolid forming, pressure die casting, injection molding or extrusion.

[0020] Also accordingly, a metal matrix composite (MMC) is provided. The MMC comprising a metal matrix of a first metal or alloy; and numerous sub-millimeter dimension embedded particles of a metal-oxide ceramic distributed throughout the metal matrix, wherein 0.005 to 10 wt.% calcium is present, and a concentration of calcium within the embedded particles and surrounding the embedded particles is more than double a concentration of the calcium in the metal matrix away from the embedded particles.

[0021] The oxides of calcium may be more highly concentrated at a periphery of the particles than within the ceramic clusters, linking the first metal and the ceramic clusters. The ceramic particles preferably include titanium dioxide (TiO₂), calcium oxide and aluminum oxide, and the first metal is aluminum or an alloy of aluminum. The ceramic particles and first metal or alloy are preferably present in a ratio of between 80:20 to 0.1:99.9 wt. %; more preferably in a ratio of between 65:35 to 1:99 wt. %, or between 55:45 to 5:95 wt. %, as specifically shown.

[0022] Furthermore a method is provided for reducing melt loss due to calcium defects in parts formed from an aluminum or aluminum alloy melt, the method comprising estimating a molar amount of calcium present, and adding at least an equal molar amount of rutile titania to the aluminum or aluminum alloy melt.

[0023] Further features of the invention will be described or will become apparent in the course of the following detailed description.

Brief Description of the Drawings

[0024] In order that the invention may be more clearly understood, embodiments thereof will now be described in detail by way of example, with reference to the accompanying drawings, in which:

[0025] FIGs. 1a,b show separation of rutile titania in molten aluminum shown on an X ray image and photograph, respectively; and

[0026] FIGs. 2a,b,c,d are images at increasing magnifications of an extracted sample of a wedge in the casting campaign, and an EDS analysis of calcium at the largest magnification.

Description of Preferred Embodiments

[0027] Herein a MMC material system is described, the material system formed of at least a metal matrix that includes aluminum, and embedded reinforcements dispersed within the matrix. The reinforcements are composed of, or coated with ceramic particles, which may be a ceramic oxide, boride, carbide, nitride or graphite. More preferably the ceramic is an oxide or boride, or a ceramic that has a naturally formed oxidization layer, such as silicon carbide, for example. More preferably the ceramic is an oxide, such as titania in a crystal form other than anatase. More preferably the ceramic is rutile titania, brookite titania, or a combination thereof. Most preferably the ceramic is rutile.

[0028] An interface region is formed at the boundaries between the ceramic and matrix. The interface region includes Ca, and the concentration of Ca in the interface region is far greater than the concentration of Ca in the metal matrix. Preferably the Ca is effectively not present in the metal matrix away from the interface region. The Ca may be effectively only in the interface region, or effectively only in the interface region and within the reinforcements. The preferred order for affinities for oxygen of these metals is preferably calcium, matrix metal and the ceramic (and its constituents). Rutile TiO₂ has a particular ability to react with calcium in the metal matrix, and thus even though calcium can be a problem in aluminum

and aluminum alloys, it can be effectively used to promote the integration of ceramics since its reaction has been found to remove it from the matrix.

[0029] A method of producing a MMC involves mixing reinforcements with an aluminum-containing molten metal, and between 0.005 and 10 wt.% Ca (more preferably 0.005 to 5 wt. %, and more preferably from 0.01 to 2.5 wt. %), wherein the reinforcements are particles that have a surface bearing at least 20% of titanium oxide (TiO_2), in a crystal form other than anatase (preferably rutile), and cooling the mixture to produce a solid metal matrix composite. The titania may include brookite, which is expected to equally improve integration, given similarities in the crystal structures of the two polymorphs. The crystal structure of brookite is compatible with rutile, and brookite can grow epitaxially on rutile. Anatase, on the other hand, has a very different crystal structure, which is evidently less compatible with the formation of the calcium-containing composition observed. It is noted that brookite is a relatively scarce polymorph of rutile.

[0030] The reinforcements may be ceramic or cermet, and may consist of ceramic compositions having a variety of grains of different composition, crystal form, or shape. The particles are typically dense, if a strong MMC is desired. Some properties of ceramics are achieved only with particles smaller than a given size, and frequently the size is in the nanometer scale. The addition of Ca, given the markedly improved integration of rutile with Al-containing metals and alloys, may allow for higher ceramic content in the MMC, or for better integration of finer rutile reinforcements, or other reinforcements coated with rutile powder.

[0031] The reinforcements typically have all dimensions smaller than 1 cm and may be nanostructured or microstructured, coated with rutile, a cermet of rutile in a metal (the same as or different than the matrix metal), or monolithic. The reinforcements may have any distribution of sizes, angularities, or surface areas, although are expected to have at least one sub-millimeter, and often sub-micron dimension. Substantially equiaxed powders may be preferable in many applications, although fibres, filaments and rods, and platelets, discs or flakes may be useful in others. The presence of rutile on the surface of the powders permits the formation of a Ca containing boundary layer that links the metal matrix and the particles which may improve adherence of the MMC, and may improve longevity of the MMC, and further attracts the Ca away from the metal matrix.

[0032] The molten metal is preferably Al or an alloy of Al (with at least 10%, or more preferably 20, 30, 40, 50, 60, 70, 80, 90, 95, 97, 99 wt.% or more of Al). If a high ceramic content is desired (i.e. more than 35 wt.%), the alloy may preferably not contain Mg. Even

moderately small amounts of Mg (2%) have been found to impair the integration of high concentrations of rutile by liquid Al, although greater amounts of Ca, and other alloys of Al may reduce this effect. The metal matrix may contain moderately small amounts of boron, or other metals, and may include other reinforcements (be they ceramic or other) not linked to the matrix, by a systematically Ca-containing boundary layer.

[0033] If a molten alloy of Al is used, preferably no alloying metal present in substantial quantities, have a higher affinity for oxygen than Ca. Any alloying metals included preferably do not react more readily with the reinforcements than Al, or otherwise impede the reactions between the Al, Ca, and ceramic.

[0034] The MMC may be composed entirely of the monolithic ceramic powder, molten metal, and Ca, each with their respective impurities. Alternatively other reinforcements, solid metals in the molten metal (forming a semi-solid) or other alloying materials, or other materials may be present, and so the mixture may be at least 60 wt.%, more preferably 80 wt.%, more preferably 90 wt.%, more preferably 95 wt.%, more preferably 97 wt.% of the powder and molten metal.

[0035] Cooling the mixture to produce a solid metal matrix composite may involve known processes such as: sandcasting, die casting, centrifugal casting, compocasting, thixocasting, rheocasting, thixomolding or other semisolid forming, pressure die casting, injection molding or extrusion.

[0036] This method may produce a metal matrix composite (MMC) formed of a metal matrix of a first metal or alloy; and numerous sub-millimeter dimension embedded particles distributed uniformly throughout the metal matrix, wherein 0.005 to 10 wt.% calcium is present, but is at least mostly confined within a boundary layer produced around the ceramic particles. For example a concentration of calcium confined to the embedded particles and surrounding the embedded particles, is more than double a concentration of the calcium in the metal matrix away from the embedded particles. The concentration of calcium within and around the embedded particles may be more than 10 times, more than 50 times, and more than 100, or 1000 times the concentration of calcium in the metal matrix away from the embedded particles.

[0037] With the formation of a boundary layer around the embedded particles, the calcium may be more highly concentrated at a periphery of the particles than within the particles themselves. The boundary layer may better link the first metal and the ceramic clusters. The embedded ceramic particles may include titanium, calcium, oxygen, and

aluminum, and the first metal may be aluminum or an alloy of aluminum, and preferably the embedded ceramic particles were prepared from compounds of known purities of rutile titanium oxide (TiO_2), with calcium oxide and substantially aluminum oxide, and the first metal is aluminum or an alloy of aluminum.

[0038] As calcium is a known impurity for Al, and as rutile titania is abundant, it also makes sense to treat the rutile as an additive that compensates for and effectively removes the Ca from Al. As such rutile titania may be used to reduce melt loss, energy, labour, and processing when an aluminum metal or alloy is known to contain calcium.

EXAMPLES

[0039] Applicant has experimented with the incorporation and integration of TiO_2 in liquid aluminum. Specifically, approximately 50 g of rutile TiO_2 powder (99.9%, $<5\ \mu\text{m}$, $4.17\ \text{g/cm}^3$, product No. 224227, Sigma-Aldrich), was folded in an aluminum foil and placed at the bottom of a steel crucible. Commercially pure aluminum ($>99.9\%$, Al P0404, AIM Metals and Alloys) was melted in an electric furnace at a temperature of approximately $720\ ^\circ\text{C}$ and then poured in the steel crucible over the foil which freed the powder as it melted. A total of 5 slugs were produced in this manner and it was observed during these trials that the TiO_2 tended to rise to the surface. An X-Ray inspection system (model Y Multiplex 5500 M, 225 kV, variofocus tube, YXLON) was used to examine the slugs and revealed the presence of large porosity in their upper portions, a typical radiograph being shown in FIG. 1a. Large defects are shown in the upper portions of the slug by the radiograph. The slugs were then sliced for internal examination, and are photographed (presented as FIG. 1b). The presence of large porosity originating from solidification shrinkage was observed as well as some TiO_2 powder clustered inside the cavities. Some white TiO_2 powder was found clustered in some of the cavities.

[0040] An examination with a scanning electron microscope (Hitachi SU-70 FEG SEM) revealed that the TiO_2 powder was mainly located in the shrinkage porosity and had remained unwetted by aluminum. The chemical composition provided by the energy dispersive X-ray spectroscopy (EDS) system (Oxford EDS INCA 300) showed the presence of aluminum, oxygen and titanium along with some contaminants.

[0041] No evidence was found of aluminum reacting with the TiO_2 . Aluminum has a very high affinity for oxygen, its reaction producing aluminum oxide, Al_2O_3 . This compound is more stable than titanium oxide, TiO_2 , and some reduction would thus be expected when TiO_2 additions are made to liquid aluminium unless kinetic barriers are present. Moreover,

titanium has limited solubility in liquid aluminium (<1 wt% at 800°C) and titanium aluminides would be expected to form even when a small amount of TiO₂ is reduced. With sufficient mass fractions of TiO₂ in liquid aluminum, aluminum oxide and titanium aluminide would be expected to be produced according to the following exothermic reaction: $3 \text{TiO}_2 + 7 \text{Al} \rightarrow 2 \text{Al}_2\text{O}_3 + 3 \text{TiAl}$. The results from the gravity casting showed a tendency for TiO₂ to agglomerate and poor integration with liquid aluminum. There is no sign of a chemical reaction between the Al and titania.

[0042] Additional tests to evaluate the incorporation of TiO₂ were carried out with the stir-casting technique and an attempt to produce wedges by high pressure die casting with this slurry was made. In these tests, anatase TiO₂ was used (≥99%, <44 μm, 3.9 g/cm³, product No. 248576, Sigma-Aldrich). Approximately 90 kg of aluminum (>99.9%, Al P0404, AIM Metals and Alloys) was melted in an electric furnace and a vortex in liquid aluminum was created by the rotating impeller of a mixer. The anatase was first heated to 300°C for at least 1 hour to remove moisture and a total of 9 kg was poured into the vortex by incremental additions of 300 g batches.

[0043] Agglomeration and lack of wetting were again observed with this mode of incorporation and once the vortex stopped, the TiO₂ immediately separated from the melt and floated to the surface. Although the supplier specified a density of 3.9 g/cm³ for the anatase TiO₂, the apparent density was measured to be 0.5 g/cm³ and combined with the lack of wetting, is believed to account for the observed rise to the surface of liquid aluminum (ρ_{Al}= 2.4 g/cm³). The high pressure die casting trials also failed to produce presentable wedges and the separation of the solid TiO₂ from the liquid aluminium was the main reason.

[0044] Experiments were performed to assess the effect on integration of two different TiO₂ forms (rutile and anatase) having different granulometry and hence different apparent densities, and the effects of small additions of boron, magnesium and calcium metals (that could modify wetting of aluminum with TiO₂). A two-level screening design comprising 16 trials was selected, having for response variable the amount of TiO₂ that could be incorporated in aluminum.

[0045] These tests were performed in a small furnace with a capacity to melt approximately 5 kg of aluminum. A mechanical stirrer (IKA, model RW20DWMNS1, Fischer Scientific) mounted with a steel impeller was used for mixing the TiO₂. Oxidation of aluminum was reduced with argon at a flow rate of 15 L/min that was supplied by a ring placed above the crucible and made with a copper tube (1/4") having perforated holes.

[0046] The anatase was the same as described above (product No. 248576, Sigma-Aldrich) while the rutile was supplied by Rio Tinto Iron and Titanium (>97% pure, UGSTM, 300-350 μm , $\rho = 3.9 \text{ g/cm}^3$, $\rho_{app} = 1.87 \text{ g/cm}^3$). In all instances, the TiO_2 powder was heated to 300°C for at least 1 hour to remove moisture. Magnesium (99.9%, Rand Alloys) was added to the melt while calcium (Al-10%Ca, Rand Alloys) and boron (Al-4%B, AIM Metals and Alloys) were added as master alloys. Magnesium, calcium and boron were weighed and added to liquid aluminum before the TiO_2 additions. As it was unknown what amount of titania would be accepted by the melt, fixed amounts of Mg 2 wt.%, Ca 2 wt.%, and B 1 wt.% with respect to the initial quantity of pure liquid aluminum were used or not for each trial.

[0047] Regardless of whether Mg, Ca, or B were included, anatase titania exhibited very poor mixing, and separated readily once the mixer stopped. In all cases, except with Mg and Ca and no B (which showed poor mixing/lumpiness), less than 13 wt.% was incorporated, and typically at around 10 wt.% it is clear that no more titania can be added. Sparking and flaring was also observed, indicating poor integration.

[0048] Rutile titania, which has exactly the same chemical composition as anatase titania, exhibited very different mixing. While differences in the apparent densities of the anatase (45 microns-0.5 g/cm^3) vs. rutile (300 to 350 microns-1.87 g/cm^3) were considered to possibly have had some effect (liquid aluminium has a density of 2.4 g/cm^3), subsequent experiments with different diameter powders and apparent densities suggest that there is another reason for the different behaviours of these powders, perhaps owing to the crystal structure itself.

[0049] With no Mg, Ca or B, the rutile titania did not rise after mixing, but large lumps were included in the melt. Adding only Mg, good mixing is observed up to about 30 wt.%, although surface sparking is observed at higher concentrations of the rutile. Adding B only, or with the Mg makes the clumping worse, and results in separation of the powder once mixing stops.

[0050] With Ca but no Mg, the rutile titania exhibited good mixing, little sparking, and no surface segregation when the mixing is stopped. Much more titania could be included. The experiments stopped at 55 wt.%. The slurry with 55 wt.% titania was thick and had a consistency similar to semisolid aluminum billets. The addition of B to this had no appreciable effect.

[0051] With Ca and Mg, the mixing was fair, and 55 wt.% of rutile was added. There were some lumps, but no segregation when mixing stopped. Inspection showed wetting was

less than without the Mg, and the mixture was not as uniform. With B in addition, there is very poor wetting, and long lived sparks during the addition of the rutile. About 37 wt.% of rutile was added.

[0052] The results of the experiments are clearly that using the rutile polymorph had a substantial positive correlation with the ability to integrate more titania in aluminum, that the inclusion of calcium had a substantial positive correlation with the ability to integrate more titania in aluminum (individually or jointly) and that the inclusion of B and Mg are jointly negatively correlated with integration of titania in molten aluminum.

[0053] Applicant then produced wedges by high pressure die casting two formulations. In the first, 35 kg of commercially pure aluminum (>99.9%, Al P0404, AIM Metals and Alloys) were melted in an electric furnace. To this, 7 kg of aluminum-calcium master alloy (Al-10%Ca, Rand Alloys) was added. The rutile (>97%, 300-350 μm , $\rho = 3.9 \text{ g/cm}^3$, $\rho_{app} = 1.87 \text{ g/cm}^3$, UGSTM, Rio Tinto Iron and Titanium) was heated to 300°C for at least 1 hour to remove moisture and mixed to the liquid aluminum in batches of around 300 g until an amount of 51 kg was added. The additions were made with the stir-casting technique using a mixer with a graphite shaft and impeller. The melt temperature was maintained at $700 \pm 10^\circ\text{C}$ during the TiO_2 additions. As in the previous tests, aluminum oxidation was reduced with argon (38 L/min) supplied by a ring made with a copper tube (1/4") and perforated holes placed above the crucible. The final composition of the mixture in weight per cent was: Al-0.75%Ca-54.8% TiO_2 and a series of 22 wedges were cast with it.

[0054] The second casting campaign was carried out with boron addition. The preparation procedure was the same as the first campaign except that the amounts of components were: 22 kg of the commercially pure aluminum, 4.4 kg of the Al-Ca master alloy, 5.5 kg of Al-B master alloy (Al-4%B, AIM Metals and Alloys) and 36.5 kg of rutile. The final composition of the mixture in weight percent was: Al-0.64%Ca-0.32%B-53.4% TiO_2 and a series of 19 wedges were cast.

[0055] A high pressure die casting press (Buhler, SC N/53) was used with a die to cast wedge plates and the intensification pressure that was typically 850 bar. Each wedge, with its feeding system and overflows, weighed approximately 2.5 kg and had the following dimensions: L = 190 mm, W = 100 mm, T = 10 to 15 mm. During the first campaign, it was observed that the slurry was thinner at the beginning and thicker towards the end and this may have caused some variations in the amount of ceramic particles in the castings. The consistency of the slurry for the second casting campaign appeared more uniform, probably because of the slightly greater depth of the mixer impeller during preparation. Although the

castings produced in both campaigns had, in some instances, surface imperfections, they were all visually in fair condition considering that no attempts were made to optimize the casting parameters.

[0056] The solidification of pure aluminum is accompanied with relatively high volume shrinkage ($\sim 6.7\%$) and this is often accompanied by hot tearing. While some modest amount of hot tearing was observed, it is believed to be possible to avoid these defects by optimizing the casting parameters. These castings were subjected to radiographic inspections and metallographic analyses that comprised optical microscopy, scanning electron microscopy (SEM), and energy dispersive X-ray spectroscopy (EDS).

[0057] The plates were examined with the X-ray inspection system, revealing the presence of plume-like zones in light gray which were less dense than the background. Since the specific gravity of TiO_2 is 3.9 and that of solid aluminum is 2.7, lighter zones are thus considered poorer in TiO_2 . The density variations may originate from the feedstock with the Al- TiO_2 mixture being not entirely uniform or from segregation produced by shear forces during mold filling.

[0058] The castings were cut longitudinally at the center. The left hand side of the plate was used to evaluate specific gravity while a sample for microscopy evaluation approximately 4 cm x 1.25 cm was extracted from the right hand side, at the mid height. Specific gravity measurements were carried out using Archimedes' principle assuming a law of mixture for pure aluminum and TiO_2 and values for their respective specific gravity of 2.7 and 3.9. Even though the values are conservative, as porosity is not accounted for, the TiO_2 contents are well below expected, suggesting that a reaction between TiO_2 and aluminum may have taken place.

[0059] Small samples taken from the right hand side of the wedges were first examined by optical microscopy from which mosaics were made. The one for casting No. 6 at the Al-0.75%Ca-54.8% TiO_2 composition is shown in FIGs. 2a,b and was found to be typical. FIG. 2a shows TiO_2 particles imbedded in aluminum and look as though they are sandwiched between a layer of aluminum at the top and bottom. This phenomenon has also been noticed with semi-solid aluminum and is mainly caused by the presence of a shearing gradient in the injected slurry which is maximal at the interface with the die. This gradient acts as a driving force for segregation. The layer is however quite thin ($\sim 1\text{mm}$) and overall, the particles seem to be relatively well wetted and distributed.

[0060] The samples were then examined at larger magnifications (FIG. 2c) with a scanning electron microscope (Hitachi SU-70 FEG SEM). FIG. 2c provides a picture of embedded ceramic particles around which bright layers with thin border lines can be seen. These layers were observed around all the embedded ceramic particles that were examined, whether boron was added or not. An analysis with an energy dispersive X-ray spectroscopy (EDS) system (Oxford EDS INCA 300) showed that most of the calcium was contained in that layer (see FIG. 2d, calcium shown in white). A series of EDS measurements were then carried out to map variations in chemical compositions. As shown in FIG. 2c, five locations that systematically corresponded to the following, were used to generate measurements identified as Spectra 1 to 5: Spectrum 1: In the aluminium matrix; Spectrum 2: In the dark layer around the particle; Spectrum 3: In a white part of the particle, just next to the dark layer; Spectrum 4: Inside the particle, at about half the radius; Spectrum 5: Inside the particle, approximately at the center. This analysis generated Table 1 data.

Table 1: EDS measured compositions

Spectrum	O	Al	Si	Ca	Ti	Fe	Total
Spectrum 1		99.69			0.31		100.00
Spectrum 2	15.32	80.47	0.67	3.18	0.37		100.00
Spectrum 3	39.34	5.61		0.40	54.08	0.56	100.00
Spectrum 4	38.59	6.74		0.31	53.68	0.68	100.00
Spectrum 5	37.45	26.49		0.18	35.88		100.00

[0061] For each sample extracted from the 6 castings, this analysis was repeated on five different particles that were randomly selected for a total of 30 measurements (15 from castings without boron (campaign No. 1) and 15 from castings with boron (campaign No. 2). Differences between the results of the two campaigns are not significant and boron was not detected due to its small content and its low atomic weight. The discussion below thus applies to both sets of results.

[0062] The composition at Spectrum 1 was taken in the matrix and consisted, as expected, almost exclusively of aluminum, with some reduced titanium. Spectrum 2, taken in

the dark layer around the particles, is rich in aluminum and oxygen but also contains a fair amount of calcium. The presence of this calcium-containing layer bordering the embedded particles provides an explanation for the positive effect that calcium additions had in promoting integration of the particles with aluminum. The titanium content is small at this location. Spectrums 3, 4 and 5, all taken in the pale portion of the particles, show the presence of titanium, oxygen and aluminum at roughly 50 wt%, 35 wt% and 15 wt%, respectively. The 15 wt% aluminum content is relatively high and suggests that a reaction between TiO_2 and aluminum took place. The weight percentages of these 3 elements correspond to a compound with an approximate stoichiometry of $\text{Ti}_2\text{O}_4\text{Al}$ or (with respect to 1 mole of atoms) $\text{Ti}_{0.286}\text{O}_{0.571}\text{Al}_{0.143}$. A brief literature review of the Ti-Al-O ternary system has not revealed that compounds with this approximate composition have been reported. Although titanium aluminides such as Ti_3Al and TiAl have some oxygen solubility, the amount measured here (~35 wt%) appears too high to conclude that they are present, but this possibility is not ruled out.

[0063] In conclusion, the preparation of an aluminum feedstock containing high concentrations of rutile TiO_2 (in excess of 30 wt%, 40 wt.% and 50 wt.%) was made possible by adding a small quantity (<0.75 wt%) of calcium in the aluminum. Boron additions (~ 0.3 wt%) were not found to have detrimental effects. Magnesium additions were also made (<2 wt%) but the effect was found to be small and negative, despite the prevalent opinion that Mg is the preferred wetting agent for aluminum. Marked differences were observed between anatase and rutile. EDS analysis showed the systematic presence of thin boundary layers around the embedded particles containing calcium. The considerable positive effect of calcium to the integration of TiO_2 was attributed to the formation of this layer. The particles which initially consisted of TiO_2 (60 wt% titanium and 40% oxygen) reacted and were found after integration to the melt to consist of titanium (50 wt%), oxygen (35 wt%) and aluminum (15 wt%).

[0064] Two test bars were tested to estimate strength. The bars were composed of a matrix of Aluminum (> 99wt% purity) with particles that were TiO_2 Rutile (> 97 wt. % purity) + Silica (< 3 wt. %) The particle granulometry was dp50 of 300-350 μm . The particle content in the matrix was ~ 55 wt. %. The plates were extracted from high pressure die cast plates in the as-cast condition (no heat treatment, tempering or annealing). The bars were finished as required by ASTM standards for strength testing. Nonetheless, useful information about the bars were observed. The Young's modulus for the material was observed to be about 80 ± 0.5 GPa; the yield strength was found to be 54 ± 2 MPa; the tensile strength was found to

be 64 ± 10 MPa; and the elongation was found to be $1.5 \pm 1\%$. These values appear to compare favourably with commercially available MMCs.

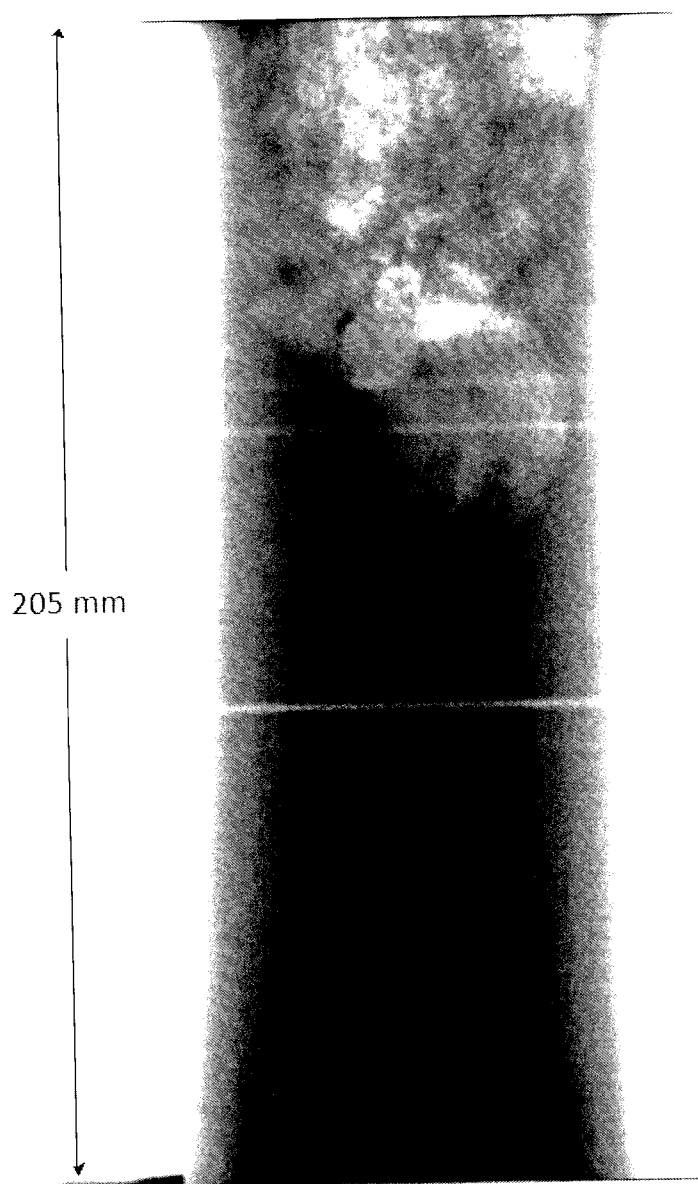
[0065] A casting campaign was carried out with finer rutile powders (> 99 wt. % purity), and found that even with nominally 30-50 μm powders, 55 wt. % of rutile could be incorporated, although this was approaching a limit for the specific composition.

[0066] Other advantages that are inherent to the structure are obvious to one skilled in the art. The embodiments are described herein illustratively and are not meant to limit the scope of the invention as claimed. Variations of the foregoing embodiments will be evident to a person of ordinary skill and are intended by the inventor to be encompassed by the following claims.

Claims:

1. A method for producing a metal matrix composite comprising:
mixing a reinforcement with an aluminum-containing molten or semisolid metal or alloy and between 0.005 and 10 wt.% calcium (Ca), wherein the reinforcement is composed of particles each having a surface bearing at least 20% of titanium oxide (TiO_2), and the TiO_2 is predominantly of crystal form other than anatase; and cooling the mixture to produce a solid metal matrix composite.
2. The method of claim 1 wherein the reinforcement is a cermet or ceramic powder including the TiO_2 , or a compound coated with the TiO_2 .
3. The method of claim 1 or 2 wherein the TiO_2 is in a rutile crystal form.
4. The method of claim 1, 2 or 3 wherein the mixture consists of at least 60 wt.%, more preferably 80 wt.%, more preferably 90 wt.%, more preferably 95 wt.%, more preferably 97 wt.% of the reinforcement and molten metal.
5. The method of any of claims 1-4 wherein the molten or semisolid metal is liquid aluminum of a predetermined purity.
6. The method of any of claims 1-4 wherein the molten metal includes aluminum, and at least one alloying metal in liquid or semisolid form with the aluminum, other than magnesium.
7. The method of any of claims 1-6 wherein the particles are spherical, cubic, prismatic, polyhedral, angular, amorphous, elongated, rod-like, tubular, conic, fibrous, filamentary, platelet-like, disc-like, irregular, or any combination of the above.
8. The method of any of claims 1-7 wherein the surfaces of the particles are flat, or curved, smooth or rough, randomly textured or patterned, concave or convex, or any combination of the above.
9. The method of any of claims 1-8 wherein the particles have a predefined distribution of dimensions, with less than 10% of the powders having dimensions greater than a maximum dimension, which is less than 1 cm, and with less than 10% of the powders having dimensions smaller than a minimum dimension, which is greater than 10 nm.
10. The method of any of claims 1-9 wherein each surface of the typical particle bears at least 20% of TiO_2 .

11. The method of claim 10 wherein each surface of the typical particle bears at least 60% of TiO_2 .
12. The method of any of claims 1-11 wherein cooling the mixture to produce a solid metal matrix composite comprises: sandcasting, die casting, centrifugal casting, compocasting, thixocasting, rheocasting, thixomolding or other semisolid casting, pressure die casting, injection molding or extrusion.
13. A product of the method of any of claims 1-12.
14. A metal matrix composite (MMC) comprising:
 - a metal matrix of a first metal or alloy; and
 - numerous sub-milimeter dimension embedded particles of a ceramic distributed throughout the metal matrix,
 - wherein 0.005 to 10 wt.% calcium is present, and a concentration of calcium within the embedded particles and surrounding the embedded particles is more than double a concentration of the calcium in the metal matrix away from the embedded particles.
15. The MMC of claim 14 wherein the oxides of calcium are more highly concentrated at a periphery of the particles than within the ceramic clusters, linking the first metal and the ceramic clusters.
16. The MMC of claim 14 wherein the ceramic particles include crystals of rutile titanium oxide (TiO_2), calcium oxide and aluminum oxide, and the first metal is aluminum or an alloy of aluminum.
17. The MMC of claim 14 wherein the ceramic particles are composed of, or coated with a ceramic oxide, boride, carbide, nitride or graphite.
18. The MMC of claim 14 wherein the ceramic particles are composed of, or coated with an oxide or boride, or a ceramic that has a naturally formed oxidization layer.
19. A method for reducing melt loss due to calcium defects in parts formed from an aluminum or aluminum alloy melt, the method comprising estimating a molar amount of calcium present, and adding at least an equal molar amount of rutile titania to the aluminum or aluminum alloy melt.



FIG, 1a

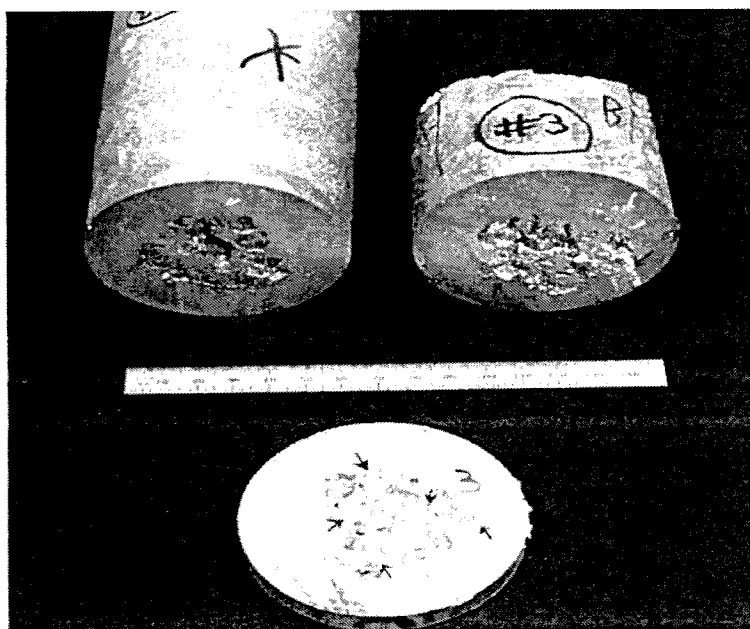
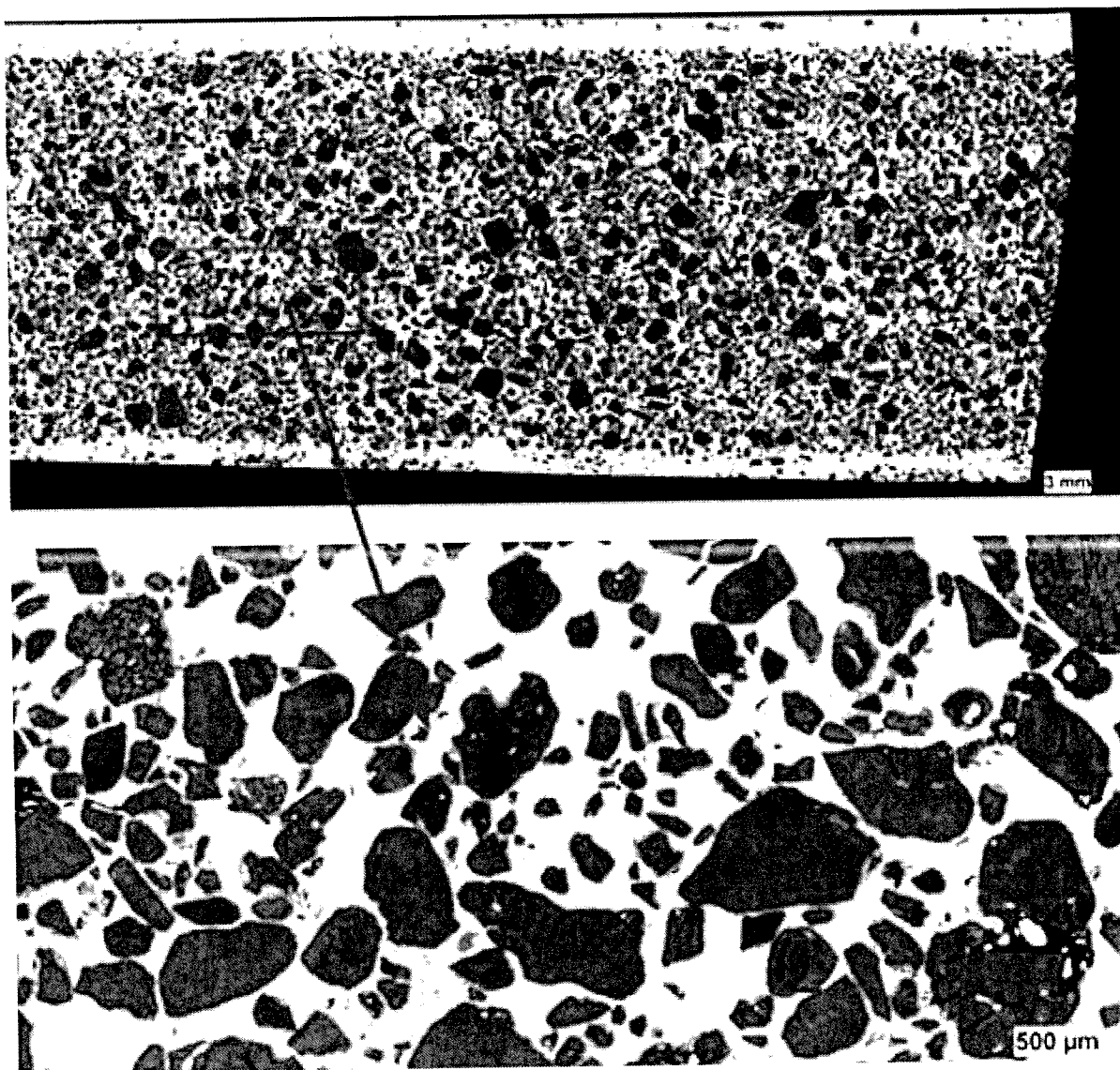
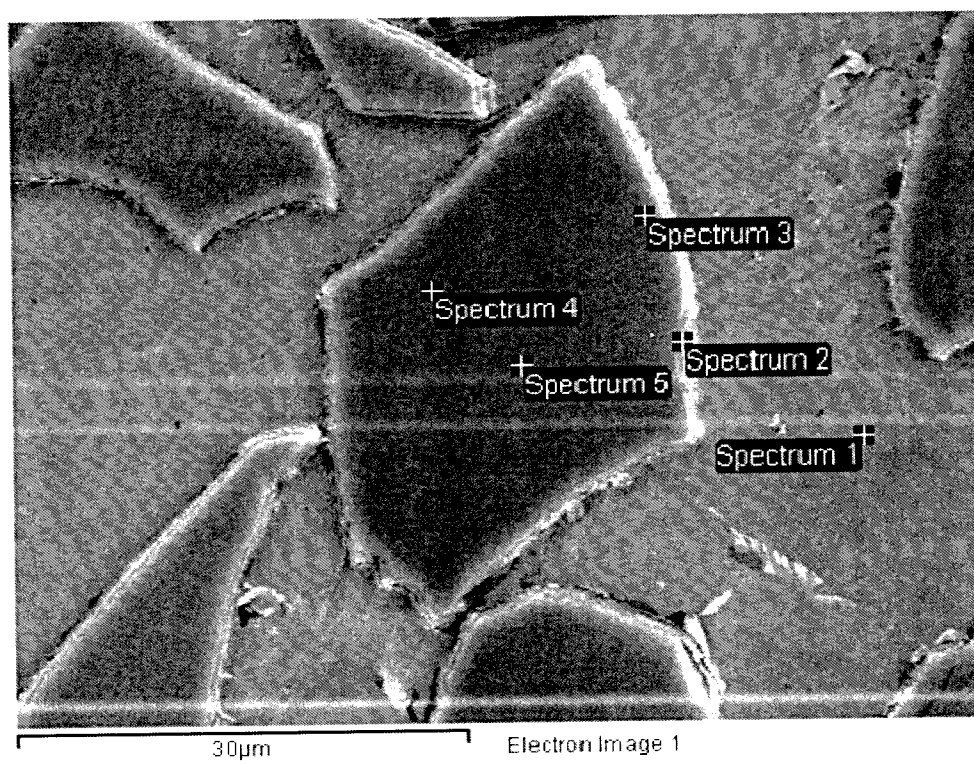


FIG. 1b

FIG. 2a**FIG. 2b**

**FIG. 2c**

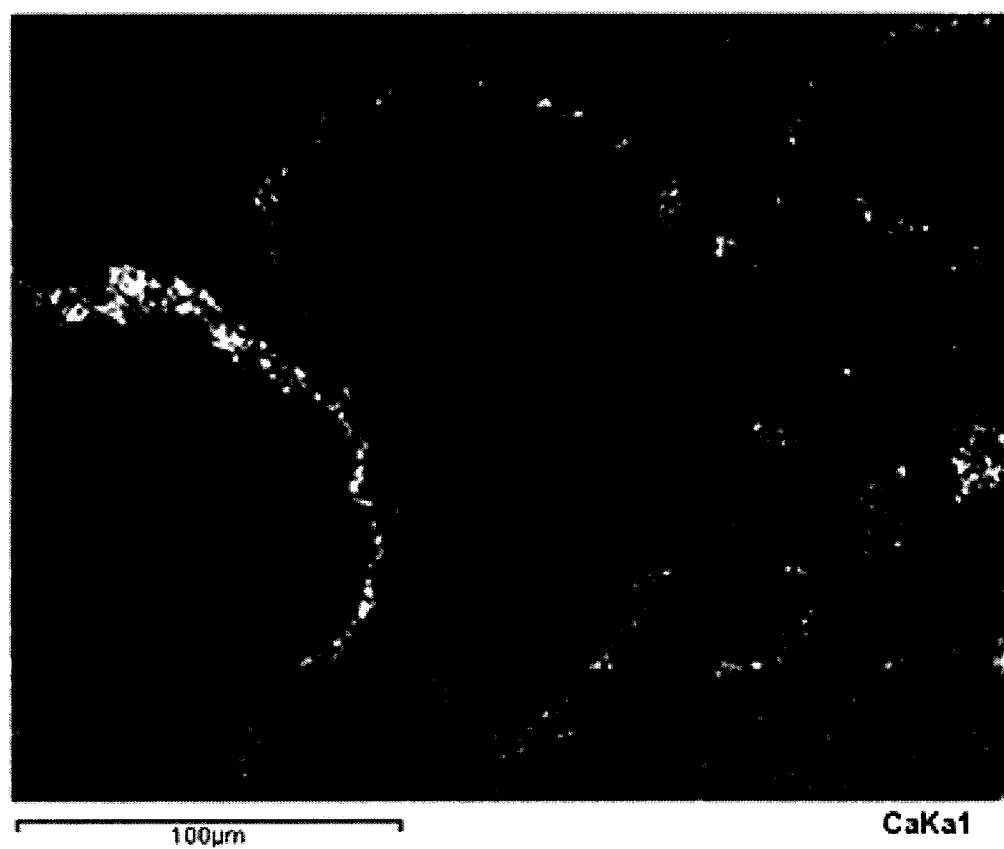


FIG. 2d

INTERNATIONAL SEARCH REPORT

International application No.

PCT/CA2014/000102

A. CLASSIFICATION OF SUBJECT MATTER

IPC: **C22C 32/00** (2006.01), **C22C 21/00** (2006.01), **C22C 29/12** (2006.01)

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC: **C22C 32/00** (2006.01), **C22C 21/00** (2006.01), **C22C 29/12** (2006.01)

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic database(s) consulted during the international search (name of database(s) and, where practicable, search terms used)

EPOQUE (epodoc, txten, alloys); Canadian Patent Database; Total Patent; Google; Science Direct; Keywords: Ti, titanium, titania, rutile, Ca, calcium, ceramic, particle, matrix, aluminium, aluminum, Al.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	JP07-102331, (TSUCHITORI I. et al.), 18 April 1995 (18-04-1995) **machine translation of abstract, [0017], [0018], [0019], [0021]**	1-8, 10-19
X	JP60-159137, (KOMURO K., et al.), 20 August 1985 (20-08-1985) **machine translation of abstract**	14, 15, 17, 18
X	JP01-108326, (MORI S. et al.), 25 April 1989 (25-04-1989) **machine translation of abstract**	14, 15, 17, 18
X	EP0324706A2, (DWIVEDI R. K. et al.), 19 July 1989 (19-07-1989) **abstract, claims, col. 3 L13-14, col.5 L18-26, examples 1-11**	14, 15, 17, 18
X	JP60-138041, (OKI T. et al.), 22 July 1985 (22-07-1985) **machine translation of abstract, Figure 3**	14, 17, 18

☐ Further documents are listed in the continuation of Box C.☒ See patent family annex.

* "A" "E" "L" "O" "P"	Special categories of cited documents: document defining the general state of the art which is not considered to be of particular relevance earlier application or patent but published on or after the international filing date document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed	"T" "X" "Y" "&"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art document member of the same patent family
--------------------------------------	--	--------------------------	--

Date of the actual completion of the international search
29 April 2014 (29-04-2014)Date of mailing of the international search report
20 May 2014 (20-05-2014)Name and mailing address of the ISA/CA
Canadian Intellectual Property Office
Place du Portage I, C114 - 1st Floor, Box PCT
50 Victoria Street
Gatineau, Quebec K1A 0C9
Facsimile No.: 001-819-953-2476

Authorized officer

Anne-Marie Garand-Sheridan (819) 953-1612

INTERNATIONAL SEARCH REPORT
Information on patent family members

International application No.
PCT/CA2014/000102

Patent Document Cited in Search Report	Publication Date	Patent Family Member(s)	Publication Date
EP0324706A2	19 July 1989 (19-07-1989)	EP0324706A2 EP0324706A3 EP0324706B1 AT94522T AU2670488A AU621072B2 AU620191B2 AU4718989A BG60294B2 BR8900067A BR8906609A CA1321055C CA2005977A1 CN1034688A DD301869A9 DE68909061D1 DE68909061T2 DK7989D0 DK7989A DK650289D0 DK650289A EG19192A EP0375613A2 EP0375613A3 FI890121A FI93945B FI93945C HUT55729A HUT52486A IE890031L IE60954B1 IL88868D0 IL88868A IL92707D0 IN171652A1 JPH02197368A JP3247363B2 JPH02258780A KR970002030B1 MX165625B MX18774A NO890013D0 NO890013A NZ227522A PH25592A PL277121A1 PL156828B1 PT89417A PT89417B RO103408B1 RU1797603C TR23959A TR24056A US4871008A US4998578A US5023259A YU5189A ZA8900184A ZA8909761A	19 July 1989 (19-07-1989) 05 September 1990 (05-09-1990) 15 September 1993 (15-09-1993) 15 October 1993 (15-10-1993) 13 July 1989 (13-07-1989) 05 March 1992 (05-03-1992) 13 February 1992 (13-02-1992) 28 June 1990 (28-06-1990) 25 April 1994 (25-04-1994) 05 September 1989 (05-09-1989) 04 September 1990 (04-09-1990) 10 August 1993 (10-08-1993) 21 June 1990 (21-06-1990) 16 August 1989 (16-08-1989) 01 June 1994 (01-06-1994) 21 October 1993 (21-10-1993) 17 February 1994 (17-02-1994) 10 January 1989 (10-01-1989) 12 July 1989 (12-07-1989) 20 December 1989 (20-12-1989) 22 June 1990 (22-06-1990) 29 September 1994 (29-09-1994) 27 June 1990 (27-06-1990) 28 November 1990 (28-11-1990) 12 July 1989 (12-07-1989) 15 March 1995 (15-03-1995) 26 June 1995 (26-06-1995) 28 June 1991 (28-06-1991) 28 July 1990 (28-07-1990) 11 July 1989 (11-07-1989) 07 September 1994 (07-09-1994) 15 August 1989 (15-08-1989) 15 November 1992 (15-11-1992) 17 September 1990 (17-09-1990) 05 December 1992 (05-12-1992) 03 August 1990 (03-08-1990) 15 January 2002 (15-01-2002) 19 October 1990 (19-10-1990) 21 February 1997 (21-02-1997) 25 November 1992 (25-11-1992) 01 November 1993 (01-11-1993) 03 January 1989 (03-01-1989) 12 June 1989 (12-06-1989) 29 January 1991 (29-01-1991) 08 August 1991 (08-08-1991) 24 July 1989 (24-07-1989) 30 April 1992 (30-04-1992) 08 February 1990 (08-02-1990) 30 September 1993 (30-09-1993) 15 March 1993 (15-03-1993) 23 February 1993 (23-02-1993) 11 January 1991 (11-01-1991) 01 March 1991 (01-03-1991) 03 October 1989 (03-10-1989) 12 March 1991 (12-03-1991) 11 June 1991 (11-06-1991) 31 October 1990 (31-10-1990) 26 September 1990 (26-09-1990) 29 August 1990 (29-08-1990)